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- (54) Title: BENZIMIDAZOLE VASCULAR DAMAGING AGENTS
- (54) Titre: AGENTS DE DEGRADATION VASCULAIRE AUX BENZIMIDAZOLES

(57) Abstract

A group of vascular damaging agents which can be used in the preparation of medicaments for the treatment of diseases involving neovascularisation are provided. These are 5(6)-substituted benzimidazole-2-carbamates of formula (I) wherein Alk is an alkyl group, X is oxygen, sulphur, sulphinyl, sulphonyl, carbonyl (CO), thiocarbonyl (CS), sulphonyloxy, NH, iminomethylene (C=NH), N-hydroxyiminomethylene, N-alkoxyiminomethylene, dialkoxymethylene, 1,3-dioxolan-2yl, 1,1-ethenyl, a group CHR3¿ or a bond, R1¿ is hydrogen, alkylaminocarbonyl or alkoxycarbonyl, R2¿ is hydrogen, alkoxycarbonyl, cyanomethyl, cyanoethyl, alkoxymethyl or acetoxymethyl. R3¿ is hydrogen, hydroxy, alkoxy or amino, A is an optionally substituted aromatic, optionally substituted heteroaromatic, optionally substituted heteroaromatic, optionally substituted heteroaromatic, optionally substituted salts, solvates and hydrates thereof. Most of the compounds of this group are novel, in particular those in which A is an aromatic or heteroaromatic ring with substituents, particularly substituents which are phosphates or alkylphosphates. The invention therefore provides both novel compounds and pharmacological compositions with compounds within the broad definition.

(57) Abrégé

La présente invention concerne un groupe d'agents de dégradation vasculaire convenant à la péparation de médicaments destinés au traitement d'affections impliquant une néovascularisation. Ces molécules sont des benzimidazole-2-carbamates à substitution 5(6) représentés par la formule générale (I) ou certains de leurs sels, solvates et hydrates admis en pharmacie. Dans cette formule, "alk" est un groupe alkyle. X est oxygène, soufre, sulphinyle, sulphonyle, carbonyle (CO), thiocarbonyle (CS), sulphonyloxy, NH, iminométhylène (C=NH), N-hydroxyiminométhylène, N-alcoxyiminométhylène, dialcoxyméthylène, 1,3-dioxolane-2yle, 1,1-éthényle, un groupe CHR3¿ ou une liaison. R1¿ est hydrogène, alkylaminocarbonyle ou alcoxycarbonyle. R2¿ est hydrogène, alcoxycarbonyle, cyanométhyle, cyanoéthyle, alcoxyméthyle ou acétoxyméthyle. R3¿ est hydrogène, hydroxy, alcoxy ou amino. A est un groupe éventuellement substitué, notamment aromatique, hétéroaromatique, hétérocycloalkyle, alkyle ou cycloalkyle. La plupart des composés de ce groupe entrent dans le cadre de l'invention, en particulier ceux dans lesquels A est un cycle aromatique ou hétéroaromatique à substituants, particulièrement des substituants qui sont phosphates ou alkylphosphates. L'invention concerne donc aussi bien les composés de l'invention que les compositions pharmaceutiques comportant des composés ressortissants de la définition au sens large.

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Description

BENZIMIDAZOLE VASCULAR DAMAGING AGENTS

This invention relates to vascular damaging agents and particularly to the use of new and known substituted benzimidazoles in the preparation of medicaments for the treatment of diseases involving neovascularisation.

Formation of new vasculature by angiogenesis is a key pathological feature of several diseases (J Folkman, New England Journal of Medicine 333, 1757-1763 (1995)). For example, for a solid tumour to grow it must develop its own blood supply upon which it depends critically for the provision of oxygen and nutrients; if this blood supply is mechanically shut off the tumour undergoes necrotic death. Neovascularisation is also a clinical feature of skin lesions in psoriasis, of the invasive pannus in the joints of rheumatoid arthritis patients and of atherosclerotic plaques. Retinal neovascularisation is pathological in macular degeneration and in diabetic retinopathy. In all these diseases reversal of neovascularisation by damaging the newly-formed vascular endothelium is expected to have a beneficial therapeutic effect.

5(6)-Substituted benzimidazole-carbamates are known and have found use as antiparasitic agents (P. J. Islip in Burgers Medicinal Chemistry (M. E. Wolff ed.), Fourth Edition, Part II, p481, (1979)). Examples of such compounds include mebendazole, fenbendazole, oxibendazole, flubendazole, albendazole, cyclobendazole, parbendazole, dribendazole, luxabendazole, and etibendazole. Their mode of action for their antiparasitic action is believed to involve selective binding to tubulin of the target parasite while having little effect due to binding tubulin of the mammalian host (Biochim. Biophys. Acta 630, 271-278, (1980)). Some of these compounds have been shown to be antimitotic for cancer cells and one particular 5(6)-substituted benzimidazole-2-carbamate, nocodazole, has therefore been studied as an anticancer agent (Cancer Research, 36, 905-916 (1976)). No effects on neovasculature have been reported for any of these compounds.

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Some structurally-unrelated compounds which bind tubulin have been shown to have anti-vascular effects when given at their maximum tolerated dose (MTD) (S. A. Hill et al. Eur. J Cancer, 29A, 1320-1324 (1993)) but other tubulin-binding agents, such as docetaxel, have no vascular-damaging activity even when administered at the MTD.

The presence of tubulin-binding properties is then not predictive for antivascular activity.

According to the present invention there is provided the use of 5(6)-substituted benzimidazole-2-carbamates for the preparation of compositions for the treatment of diseases involving angiogenesis in which the 5(6)-substituted benzimidazole carbamate has the formula

I

15 wherein

Alk is an alkyl group

X is oxygen, sulphur, sulphinyl, sulphonyl, carbonyl (CO), thiocarbonyl (CS), sulphonyloxy, NH, iminomethylene (C=NH), N-hydroxyiminomethylene, N-alkoxyiminomethylene, dialkoxymethylene, 1,3-dioxolan-2yl, 1,1-ethenyl, a group CHR³ or a bond

 R^1 is hydrogen, alkylaminocarbonyl or alkoxycarbonyl R^2 is hydrogen, alkoxycarbonyl, cyanomethyl, cyanoethyl, alkoxymethyl or acetoxymethyl.

R³ is hydrogen, hydroxy, alkoxy or amino
A is an optionally substituted aromatic, optionally substituted heteroaromatic,
optionally substituted heterocycloalkyl, optionally substituted alkyl or optionally
substituted cycloalkyl group

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and the pharmaceutically acceptable salts, solvates and hydrates thereof.

Particular substituents that may be present on the group A include one or more substituents selected from a group Y, optionally substituted alkyl, (where substituents on such alkyl group may include one or more selected from hydroxy, amino, alkylamino, dialkylamino, halogen, carboxyl, SO₃H, sulphate, phosphate, alkoxycarbonyl, aralkoxycarbonyl, alkoxycarbonylamino, aminoalkylaminocarbonyl and cyano), halogen, hydroxy, amino, alkoxy, alkylthio, cyano, nitro, sulphate, isothiocyanate, aryl, heteroaryl and heterocycloalkyl.

Y is a group selected from phosphate, alkylphosphate, C(O)R⁴, OC(O)R⁴, SO₂R⁴, NHC(O)R⁴, NR⁵C(O)R⁴, SR⁴, S(O)R⁴, OSO₂R⁴, NHSO₂R⁴, NR⁵SO₂R⁴, SO₃H, CO₂H and CO₂R⁵ where R⁴ is a group selected from hydrogen, R⁵, OR⁵, NHR⁵, NR⁵R⁶, aryl, heteroaryl or heterocycloalkyl groups being optionally substituted with one or more substituents selected from alkyl, heterocycloalkyl, haloalkyl, hydroxy, nitro, cyano, amino, alkylamino, dialkylamino, halogen, carboxyl, SO₃H, sulphate and phosphate. R⁵ and R⁶, which may be the same or different, are each an alkyl group optionally substituted with one or more substituents selected from hydroxy, amino, alkylamino, dialkylamino, guanidino,

20 halogen, carboxyl, SO₃H, sulphate, phosphate, aryl and heteroaryl.

Some of the compounds usable in the invention are known, for example the following compounds within the following formula:

A N H O Me

These compounds are

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X Α Fenbendazole S Ph Mebendazole CO Ph Albendazole S nPr Oxibendazole 0 nPr CO Nocodazole 2-thienyl

Certain of these compounds are novel. In one embodiment the novel compounds are those of formula I in which at least one of the substituents on the group A is a group Y where Y is as hereinbefore defined. Particularly preferred are compounds defined by the formula

10 wherein

alk is an alkyl group

B is an aromatic or heteroaromatic ring

X is oxygen, sulphur, sulphinyl, sulphonyl, carbonyl (CO), thiocarbonyl (CS),

sulphonyloxy, NH, iminomethylene (C=NH), N-hydroxyiminomethylene, N-alkoxyiminomethylene, dialkoxymethylene, 1,3-dioxolan-2yl, 1,1-ethenyl, a group CHR³ or a bond

R1 is hydrogen, alkylaminocarbonyl or alkoxycarbonyl

R² is hydrogen, alkoxycarbonyl, cyanomethyl, cyanoethyl, alkoxymethyl or

20 acetoxymethyl

R³ is hydrogen, hydroxy, alkoxy or amino

Y is as hereinbefore defined

R⁷ and R⁸ are each independently H, alkyl, halogen, hydroxy, amino, alkylamino, dialkylamino, alkoxy, alkylthio, cyano, nitro, or trifluoromethyl

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with the proviso that Y is not NHC(O)Me and when B is a thiophene ring then Y is not C(O)CF₃ and when B is a 5(6)-benzimidazole ring then Y is not NHCO₂Me or NHCO₂Et

5 and the pharmaceutically acceptable salts, solvates, hydrates and prodrugs thereof.

As used herein the term "alkyl", alone or in combinations, means a straight or branched-chain alkyl group containing from one to seven, preferably a maximum of four, carbon atoms such as methyl, ethyl, propyl, isopropyl, butyl, sec-butyl, t-butyl and pentyl. Examples of alkoxy groups are methoxy, ethoxy, propoxy, isopropoxy, butoxy and t-butoxy. The term "halogen" means fluorine, chlorine, bromine or iodine.

The term "aryl" as used herein unless otherwise stated includes reference to a C₆₋₁₀ aryl group which may, if desired, carry one or more substituents selected from halogeno, alkyl, haloalkyl, alkoxy, hydroxy, amino, nitro and cyano. The term "aralkoxy" means an alkoxy group substituted with an aryl group.

The term heteroaryl is defined herein as a mono- or bi-cyclic aromatic group containing one to four heteroatoms selected in any combination from N, S or O atoms and a maximum of 9 carbon atoms. Examples of heteroaryl groups include pyridyl, pyrimidyl, furyl, thienyl, pyrrolyl, pyrazolyl, indolyl, benzofuryl, benzothianyl, benzothiazolyl, isoxazolyl, thiazolyl, isothiazolyl, imidazolyl, triazolyl, quinolyl and isoquinolyl groups.

- 25 The term heterocycloalkyl includes heterocycloalkyl groups containing 3-6 carbon atoms and one or two oxygen, sulphur or nitrogen atoms. Particular examples of such groups include azetidinyl, pyrrolidinyl, piperidinyl, homopiperidinyl, piperazinyl, homopiperazinyl, morpholinyl or thiomorpholinyl groups.
- 30 The term cycloalkyl means a cycloaliphatic group containing 3-10 carbon atoms such as, for example, cyclopropyl, cyclobutyl, cyclopentyl or cyclohexyl.

One particularly preferred group of compounds are those of formula II in which Y is a phosphate group.

Another particularly preferred group of compounds are those of formula II in which Y is a group NR⁵C(O)R⁴, R⁵ is hydrogen and R⁴ is a 1-aminoalkyl group which can be further substituted for example by a hydroxy group.

Where one or more functional groups in compounds of formula I or II are sufficiently basic or acidic the formation of salts is possible. Suitable salts include pharmaceutically acceptable salts for example acid addition salts including hydrochlorides, hydrobromides, phosphates, sulphates, hydrogen sulphates, alkylsulphonates, arylsulphonates, acetates, benzoates, citrates, maleates, fumarates, succinates, lactates and tartrates, salts derived from inorganic bases including alkali metal salts such as sodium or potassium salts, alkaline earth metal salts such as magnesium or calcium salts, and salts derived from organic amines such as morpholine, piperidine or dimethylamine salts.

Those skilled in the art will recognise that compounds of formulae I and II may exist as stereoisomers and/or geometrical isomers and accordingly the present invention includes all such isomers and mixtures thereof. The substituted benzimidazole group is capable of existing in tautomeric forms and the formulae I and II are intended to represent all tautomeric forms and the substituent AX- is in the 5(6) position.

Compounds of Formula I or II may be prepared by any process known to a person skilled in the art. Compounds of Formula I or II may be prepared by a number of processes as generally described below and more specifically in the Examples hereinafter. In the following process description, the symbols A, X and alk when used in the formulae depicted are to be understood to represent those groups described above in relation to formula I or II unless otherwise indicated. In the schemes described below it may be necessary to employ protecting groups which are then

removed during the final stages of the synthesis. The appropriate use of such protecting groups and processes for their removal will be readily apparent to those skilled in the art.

Thus according to a further aspect of the invention compounds of formulae I and II in which R¹ and R² are hydrogen may be prepared by treatment of a diamine of formula III with a 1,3-bis(alkoxycarbonyl)- S-alkyl isothiourea, for example 1,3-bis(methoxycarbonyl)-S-methyl isothiourea or

1,3-bis(ethoxycarbonyl)-S-methyl isothiourea, in a solvent such as an alcohol, for example methanol or ethanol, optionally mixed with water, at from about room temperature to the reflux temperature of the solvent for about 5 minutes to 6 hours. The reaction medium is preferably made acidic by the addition of for example an organic acid such as acetic acid or p-toluenesulphonic acid.

Compounds of formula III are either known or can be prepared by conventional procedures.

Compounds of formulae I and II can also be prepared from other compounds of formulae I and II by chemical modification. Example of such chemical modifications that may be applied are standard alkylation, acylation, reduction, oxidation, sulphation, aromatic halogenation, aromatic nitration, phosphorylation, hydrolysis, condensation, cleavage and coupling reactions. These reactions may be used to add new substituents, to modify existing substituents or to modify other parts of the molecule.

Thus for example a compound of formula I or II in which R^1 is hydrogen can be converted into the corresponding compounds where R^1 is alkylaminocarbonyl by

treatment with an alkyl isocyanate in a solvent such as tetrahydrofuran at a temperature in the range 0° to 40°C, typically room temperature.

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In another general example a thioether group in a compound of formula I or II can be converted into a sulphinyl group by treatment with periodate in an aqueous alcohol solvent such as aqueous methanol or in aqueous acetonitrile at about -20° to 50°C, for about 1 to 16 h. Alternatively this conversion can be effected by treatment with one equivalent of a peracid such as 3-chloroperbenzoic acid in a chlorinated solvent such as dichloromethane or chloroform, at a temperature of about -30°C to room temperature.

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In a further general example a thioether group in a compound of formula I or II can be converted into a sulphonyl group by treatment with two or more equivalents of a peracid such as 3-chloroperbenzoic acid in a chlorinated solvent such as dichloromethane or chloroform, at a temperature of about -30°C to room temperature.

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In a further general example a keto group in a compound of formula I or II can be converted into a carbinol group by treatment with a reducing agent for example a hydride such as lithium aluminium hydride in an ether solvent such as diethyl ether or tetrahydrofuran at a temperature of from about 0° to the reflux temperature of the solvent

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20 solvent.

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In a further general example a keto group in a compound of formula I or II can be converted into an imine by treatment with ammonia in an alcoholic solvent such as ethanol at around room temperature for an extended period, for example three weeks.

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In a further general example a keto group in a compound of formula I or II can be converted into an oxime by treatment with hydroxylamine in an alcoholic solvent such as ethanol at around room temperature to around the reflux temperature of the solvent.

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In a further general example a compound of formula I or II containing a hydroxy group can be converted into the corresponding dihydrogenphosphate ester by treatment with

for example di-tert-butyl diethylphosphoramidite in the presence of a suitable catalyst for example tetrazole in a solvent such as an ether solvent for example tetrahydrofuran at a temperature in the range -40 to 40°C, conveniently at or near room temperature, followed by treatment with an oxidising agent for example 3-chloroperoxy benzoic acid or magnesium monoperoxyphthalate at a temperature in the range -78°C to 40°C preferably -65 to -10°C. The resulting intermediate phosphate triester is treated with an acid for example trifluoroacetic acid in a solvent such as a chlorinated solvent e.g. dichloromethane at a temperature in the range -30 to 40°C conveniently at or near 0°C to give the compound of formula I or II containing a dihydrogenphosphate ester.

In another general example an O-alkyl group may be cleaved to the corresponding alcohol (OH) by reaction with boron tribromide in a solvent such as a chlorinated solvent e.g. dichloromethane at a low temperature e.g. around -78°C.

In a further general example compounds of formula I or II may be alkylated by reaction with a suitable alkylating agent such as an alkyl halide, an alkyl toluenesulphonate, an alkyl methanesulphonate or an alkyl triflate. The alkylation reaction can be carried out in the presence of a base for example an inorganic base such as a carbonate e.g. caesium or potassium carbonate, a hydride such as sodium hydride or an alkoxide such as potassium t-butoxide in a suitable solvent such as an aprotic solvent e.g. dimethylformamide or an ether solvent such as tetrahydrofuran at a temperature of around -10 to 80°C.

In a further general example compounds of formula I or II containing an amine group may be acylated by treatment with a carboxylic acid and a coupling reagent, for example dicyclohexylcarbodiimide, in a sutable solvent for example an aprotic solvent such as dimethylformamide, an ether solvent such as tetrahydrofuran, a chlorinated solvent for example dichloromethane or a solvent mixture at a temperature in the range 0° to 60°, preferably about room temperature.

In a further general example a compound of formula I or II containing an OH group can be converted into a carbamate by reaction with an alkyl isocyanate or a carbamoyl chloride in an aprotic solvent such as dimethylformamide, an ether solvent such as tetrahydrofuran, a chlorinated solvent for example dichloromethane or a solvent mixture in the presence of a base such as a tertiary amine base for example triethylamine at a temperature in the range -20° to the reflux temperature of the solvent, conveniently at or around room temperature.

In a further general example a compound of formula I or II containing an amino group can be converted into a urea by reaction with an isocyanate or a carbamoyl chloride in an aprotic solvent such as dimethylformamide, an ether solvent such as tetrahydrofuran, a chlorinated solvent for example dichloromethane or a solvent mixture in the presence of a base such as a tertiary amine base for example triethylamine at a temperature in the range -20°C to the reflux temperature of the solvent, conveniently at or around room temperature.

In a further general example a compound of formula I or II containing a hydroxy group can be converted into a carbonate by reaction with an chloroformate in an aprotic solvent such as dimethylformamide, an ether solvent such as tetrahydrofuran, a chlorinated solvent for example dichloromethane or a solvent mixture in the presence of a base such as a tertiary amine base for example triethylamine at a temperature in the range -20°C to the reflux temperature of the solvent, preferably at or around 0°C.

- Preparation of a compound of formula I or II as a single enantiomer or, where appropriate, diastereomer may be effected by synthesis from an enantiomerically pure starting material or intermediate or by resolution of the final product in a conventional manner.
- Acid addition salts of the compounds of formula I or II are prepared in a conventional manner by treating a solution or suspension of the free base I or II with about one

equivalent of a pharmaceutically acceptable acid. Salts of compounds of formula I or II derived from inorganic or organic bases are prepared in a conventional manner by treating a solution or suspension of the free acid I or II with about one equivalent of a pharmaceutically acceptable organic or inorganic base. Alternatively both acid addition salts and salts derived from bases may be prepared by treatment of the parent compound with the appropriate ion-exchange resin in a standard fashion. Conventional concentration and recrystallisation techniques are employed in isolating the salts.

Compounds according to the invention are able to destroy vasculature that has been newly formed, for example tumour vasculature, while leaving unaffected normal, mature vasculature. The ability of the compounds to act in this way may be determined by the tests described hereinafter.

The compounds according to the invention are thus of particular use in the prophylaxis and treatment of cancers involving solid tumours and in the prophylaxis and treatment of diseases where inappropriate angiogenesis occurs such as diabetic retinopathy, psoriasis, rheumatoid arthritis, atherosclerosis and macular degeneration.

The compounds of the invention may be administered as a sole therapy or in combination with other treatments. For the treatment of solid tumours compounds of the invention may be administered in combination with radiotherapy or in combination with other anti-tumour substances for example those selected from mitotic inhibitors, for example vinblastine, paclitaxel and docetaxel; alkylating agents, for example cisplatin, carboplatin and cyclophosphamide; antimetabolites, for example 5-fluorouracil, cytosine arabinoside and hydroxyurea; intercalating agents for example adriamycin and bleomycin; enzymes, for example aspariginase; topoisomerase inhibitors for example etoposide, topotecan and irinotecan; thymidylate synthase inhibitors for example raltitrexed; biological response modifiers for example interferon; antibodies for example edrecolomab and antibodies against the EGFr, HER2 receptor or VEGF receptor: and anti-hormones for example tamoxifen. Such combination

treatment may involve simultaneous or sequential application of the individual components of the treatment.

For the prophylaxis and treatment of disease the compounds according to the invention may be administered as pharmaceutical compositions selected with regard to the intended route of administration and standard pharmaceutical practice. Such pharmaceutical compositions may take a form suitable for oral, buccal, nasal, topical, rectal or parenteral administration and may be prepared in a conventional manner using conventional excipients. For example for oral administration the pharmaceutical compositions may take the form of tablets or capsules. For nasal administration or administration by inhalation the compounds may be conveniently delivered as a powder or in solution. Topical administration may be as an ointment or cream and rectal administration may be as a suppository. For parenteral injection (including intravenous, subcutaneous, intramuscular, intravascular or infusion) the composition may take the form of, for example, a sterile solution, suspension or emulsion.

The dose of a compound of the invention required for the prophylaxis or treatment of a particular condition will vary depending on the compound chosen, the route of administration, the form and severity of the condition and whether the compound is to be administered alone or in combination with another drug. Thus the precise dose will be determined by the administering physician but in general daily dosages may be in the range 0.001 to 100mg/kg preferably 0.1 to 50mg/kg.

BIOLOGICAL ACTIVITY

The following test was used to demonstrate the activity and selectivity of compounds according to the invention.

Activity against tumour vasculature measured by fluorescent dye.

The following experiment demonstrates the ability of the compounds to damage tumour vasculature.

Tumour functional vascular volume in CaNT tumour-bearing mice was measured using the fluorescent dye Hoechst 33342 according to the method of Smith et al (Brit J Cancer 57, 247-253, 1988). The fluorescent dye was dissolved in saline at 6.25 mg/ml and injected intravenously at 10 mg/kg 6 hours or 24 hours after intraperitoneal drug treatment. One minute later, animals were killed and tumours excised and frozen; 10 µm sections were cut at 3 different levels and observed under UV illumination using an Olympus microscope equipped with epifluorescence. Blood vessels were identified by their fluorescent outlines and vascular volume was quantified using a point scoring system based on that described by Chalkley, (J Natl Cancer Inst, 4, 47-53, 1943). All estimates were based on counting a minimum of 100 fields from sections cut at the 3 different levels. Results are expressed as percentage reduction in vascular volume compared to control.

The activity of compounds of the invention in this assay is shown in Table 1.

Table 1: Reduction in tumour vascular volume measured by fluorescent dye

Compound	Dose (mg/kg) (i.p.)	Time (h)	% Reduction in vascular volume
Fenbendazole	500	6	44
Mebendazole	500	6	56
Albendazole	500	6	51
Oxibendazole	100	6	43
Nocodazole	100	24	23
Compound of Example:			
1	500	6	81
2	500	6	80 `
3	50	24	22
4	50	24	53
5	50	24	99
12	50	24	51
14	50	24	39

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5				14			
		15	50	24	61		
		16	50	24	56		
40		17	50	24	62		
10		18	50	24	47		
		19	50	24	88		
		20	50	24	69		
		21	50	24	74		
15							
		The following no	on-limiting Examples illu	strate the ir	vention:		
20	5	Example 1					
		Methyl [5(6)-(4-	hydroxybenzoyl)-1 <i>H</i> -be	nzimidazol-	2yl]carbamat	te.	
		A solution of 3,4	-diamino-4`-hydroxyber	zophenone	(49.6mg, 0.2	21mmol) and 1,3-	
05		bis(methoxycarb	onyl)-S-methyl isothiour	rea (98mg, 0	0.44mmol) in	ethanol (2.5ml) was	5
25		treated with p-to	luenesulphonic acid (7m	g) and the r	nixture heate	ed at reflux for 10	
	10	minutes. The mix	ture was cooled and the	precipitate	collected by	filtration and wash	ed
		with ethanol and	hexane to give the title	compound (16mg) as a v	white solid m.p.	

Example 2

N 13.36.

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Methyl [5(6)-(4-phosphonooxybenzoyl)-1H-benzimidazol-2yl]carbamate.

A solution of methyl [5-(4-(di-tert-butylphosphonooxy)benzoyl)-1H-benzimidazol-

>258°C, ¹H-NMR (400 MHz, d6-DMSO) δ 7.80 (s, 1H), 7.65 (d, 2H, J=8Hz), 7.50

(s, 1H), 7.49 (s, 1H), 6.89 (d, 2H, J=8Hz), 3.77 (s, 3H) ppm. m/e 311 (M+). Anal. Calculated for C16H13N3O4: C, 61.73; H, 4.21; N 13.49. Found: C, 61.68; H, 4.18;

2yl]carbamate (1.5g, 3.0mmol) in dichloromethane (55ml) was cooled in an ice bath and treated with trifluoroacetic acid (6ml) dropwise. The mixture was allowed to warm to room temperature and stirred for 1 hour before solvents were removed under reduced pressure. The residue was triturated with ether to give the title compound (1.13g) as a white solid m.p. >258°C, ¹H-NMR (300 MHz, d6-DMSO) δ: 7.84 (s,

25 1H), 7.75 (d, 1H, J=9Hz), 7.57 (d, 1H, J=8Hz), 7.52 (d, J=8Hz), 7.33 (d, 1H, J=9Hz), 3.78 (s, 3H) ppm.

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The methyl [5(6)-(4-(di-tert-butylphosphonooxy)benzoyl)-1H-benzimidazol-2yl]carbamate used as starting material was prepared as follows:

A solution of methyl [5(6)-(4-hydroxybenzoyl)-1H-benzimidazol-2yl]carbamate (100mg, 0.3mmol) in anhydrous tetrahydrofuran (1ml) stirred under a nitrogen atmosphere was treated with di-tert-butyl diethylphosphoramidite (74mg, 0.29mmol) and 1H-tetrazole (54mg, 0.78mmol) and the mixture stirred until the reaction was shown to be complete by TLC (about 1h). The cooled (-40°C) mixture was treated with 3-chloroperbenzoic acid (79mg, 0.39mmol) in dichloromethane (1ml) and stirred 10minutes before being allowed to warm to room temperature. The mixture was washed with saturated aqueous sodium bicarbonate followed by brine and the dried (MgSO4) organic phase concentrated under reduced pressure. The residue was chromatographed on silica gel eluting first with 4% methanol/dichloromethane then with 5% methanol/dichloromethane. Methyl [5(6)-(4-(di-tert-butylphosphonooxy)benzoyl)-1H-benzimidazol-2yl]carbamate (35mg) was obtained as a white solid m.p. 99-101°C.

Example 3

Methyl [5(6)-(4-phosphonooxyphenylthio)-1*H*-benzimidazol-2yl]carbamate

A solution of methyl [5(6)-(4-(di-tert-butylphosphonooxy)phenylthio)-1*H*-benzimidazol-2yl]carbamate (180mg) in anhydrous dichloromethane (10ml) at 4°C was treated with trifluoroacetic acid (1ml) and stirred for 1.5h. The mixture was allowed to warm to room temperature and concentrated under reduced pressure. Ethyl acetate was added and the mixture concentrated again. The residue was triturated with diethyl ether, washed with diethyl ether followed by acetone/water 9:1 and dried to give the title compound (134mg) as a white solid m.p. 190-193°C. Anal. Calculated for C₁₅H₁₄N₃O₆PS.3H₂O: C, 40.1; H, 4.5; N 9.3. Found: C, 40.1; H, 3.9; N 9.2. The methyl [5(6)-(4-(di-tert-butylphosphonooxy)phenylthio)-1*H*-benzimidazol-2yl]carbamate used as starting material was prepared as follows:

A solution of methyl [5-(4-hydroxyphenylthio)-1*H*-benzimidazol-2yl]carbamate

(200mg) in a mixture of anhydrous dimethylformamide (2ml) and anhydrous

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tetrahydrofuran (2ml) was treated with di-tert-butyl diethylphosphoramidite (350mg) and the mixture stirred for 48h at room temperature. The mixture was cooled to -65°C and treated gradually with magnesium monoperoxyphthalate (850mg) so that the temperature remained below -50°C. A saturated aqueous solution of sodium

- bicarbonate was added, keeping the temperature below -40°C during the addition then allowing the mixture to warm to room temperature. The mixture was extracted with three portions (50ml each) of ethyl acetate and the combined extracts washed with brine (50ml), dried (MgSO4) and concentrated under reduced pressure. The residue was purified by radial chromatography on silica gel eluting with
- dichloromethane/methanol 9:1 to give methyl [5(6)-(4-(di-tert-butylphosphonooxy)phenylthio)-1H-benzimidazol-2yl]carbamate (180mg)as a white foam.

Example 4

Methyl [5(6)-(4-aminophenylthio)-1H-benzimidazol-2yl]carbamate
 Methyl [5(6)-(4-(acetylamino)phenylthio)-1H-benzimidazol-2yl]carbamate (602 mg, 1.78 mmol) was dissolved in mixture of methanol (24 ml) and hydrochloric acid (10%, 6 ml) and heated under reflux for 16h. The solution was neutralised with ammonia solution and the methanol removed under reduced pressure. The white precipitate was
 collected by filtration, washed with water and dried in vacuo to give 392 mg of a pale yellow solid m.p. 282-284°C. m/e 298 (M*).

Example 5

Methyl [5(6)-(4-alanylaminophenylthio)-1H-benzimidazol-2yl]carbamate

A suspension of methyl [5(6)-(4-(Nα-tert-butoxycarbonylalanylamino)phenylthio)-1H-benzimidazol-2yl]carbamate (250mg) in dichloromethane (20ml) was treated with trifluoroacetic acid (4ml). The mixture was allowed to warm to room temperature and concentrated under reduced pressure. Ethyl acetate was added and the mixture concentrated again. The residue was triturated with diethyl ether to afford the trifluoroacetic acid salt of the title compound (105mg) as a white solid m.p. 178-

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182°C. m/e 485 (M+). Anal. Calculated for C₁₈H₁₉N₅O₃S.2C₂HF₃O₂ C; 43.1, H; 3.5, N; 11.4 Found C; 42.8, H; 3.8, N; 11.3

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The methyl [5(6)-(4-(N α -tert-butoxycarbonylalanylamino)phenylthio)-1H-benzimidazol-2yl]carbamate used as starting material was prepared as follows:

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A suspension of methyl [5(6)-(4-aminophenylthio)-1*H*-benzimidazol-2yl]carbamate (150mg) in anhydrous tetrahydrofuran (4ml) was treated with N-tert-butoxycarbonylalanine (100mg), cooled to -35°C and treated with 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide (100mg). The mixture was allowed to warm to room temperature and stir for 16h. Water (40ml) and ethyl acetate (20ml)

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were added and the mixture extracted with four portions of ethyl acetate (50ml each). The combined extracts were washed with brine (50ml), dried (MgSO4) and concentrated under reduced pressure. The residue was purified on silica gel eluting with ethyl acetate to give methyl [5(6)-(4-(Nα-tert-

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butoxycarbonylalanylamino)phenylthio)-1*H*-benzimidazol-2yl]carbamate (258mg) as a white solid m.p. 222-224°C. m/e 485 (M⁺).

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Example 6

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Methyl [5(6)-(4-(methoxycarbonylamino)phenylthio)-1*H*-benzimidazol-2yl]carbamate
Methyl [5(6)-(4-aminophenylthio)-1*H*-benzimidazol-2yl]carbamate(150 mg, 048
mmol) was dissolved in dimethylformamide (2 ml) and tetrahydrofuran (2 ml) and
methyl chloroformate (40 mg, 0.42 mmol) added together with triethylamine (43 mg,
0.42 mmol). The solution was stirred at 20°C for 72 h and then evaporated to dryness
under reduced pressure. The residue was purified by chromatography on silica gel
eluting with 5 % methanol/dichloromethane to give the title compound as a white

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25 solid; mp >350°C (dec.).

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Example 7

<u>Methyl [5(6)-(4-(phenylaminocarbonylamino)phenylthio)-1*H*-benzimidazol-2yl]carbamate</u>

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Methyl [5(6)-(4-aminophenylthio)-1*H*-benzimidazol-2yl]carbamate (150 mg, 0.48 mmol) was dissolved in dimethylformamide (2 ml) and tetrahydrofuran (2 ml) and

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phenylisocyanate (56.8 mg, 0.48 mmol) added together with triethylamine (50 mg, ca.0.5 mmol). The solution was stirred at 20°C for 12 h and the solvents removed in vacuo. The residue was purified on silica (ethyl acetate/hexane, 2:1) to give the title compound as a white solid; mp 335-340°C (dec.).m/e 433 (M⁺).

Example 8

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Methyl [5(6)-(4-(methoxycarbonyloxy)benzoyl)-1*H*-benzimidazol-2yl]carbamate
Methyl [5(6)-(4-hydroxybenzoyl)-1*H*-benzimidazol-2yl]carbamate (110 mg, 0.35 mmol) was dissolved in dimethylformamide (3.5 ml) and triethylamine (0.5 ml). The solution was cooled to 0°C and methyl chloroformate (50 mg, 0.52 mmol) added with stirring. The solution was stirred for 0.5 h at 0°C and the 1 h at 20°C and evaporated to dryness. The residue was dissolved in ethyl acetate (50 ml) and washed with sodium bicarbonate (sat., aq., 50 ml) and brine (50 ml), dried and evaporated. The residue was purified by radial chromatography on silica gel eluting with ethyl acetate/hexane, 1:1 followed by ethyl acetate to give the title compound as a white solid; mp 224–226°C (dec).m/e 369 (M⁺).

Prepared in an analogous fashion to Example 1 were:

20 Example 9

Methyl [5(6)-(2-methoxycarbonylphenylthio)-1*H*-benzimidazol-2yl]carbamate from methyl 2-(3,4-diaminophenylthio)benzoate (1.5g) and 1,3-bis(methoxycarbonyl)-S-methyl isothiourea (2.25g) there was obtained the title compound (1.51g) as a white solid m.p. 228-230 m/e 357 (M+). Anal. Calculated for C₁₇H₁₅N₃O₄S: C, 57.13; H, 4.23; N 11.75. Found: C, 57.39; H, 4.13; N 11.81.

Example 10

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Methyl [5(6)-(3-methoxycarbonylphenylthio)-1*H*-benzimidazol-2yl]carbamate from methyl 3-(3,4-diaminophenylthio)benzoate (246mg) and 1,3-bis(methoxycarbonyl)-S-methyl isothiourea (371mg) there was obtained the title compound (183mg) as an off-

white solid m.p. 226-228 m/e 357 (M+). Anal. Calculated for $C_{17}H_{15}N_3O_4S$: C, 57.13; H, 4.23; N 11.75. Found: C, 56.58; H, 4.31; N 11.86.

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Example 11

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Methyl [5(6)-(4-methoxycarbonylphenylthio)-1H-benzimidazol-2yl]carbamate from methyl 4-(3,4-diaminophenylthio)benzoate (830mg) and 1,3-bis(methoxycarbonyl)-S-methyl isothiourea (1.25g) there was obtained the title compound (694mg) as an off-white solid m.p. 280-282 m/e 357 (M+). Anal. Calculated for C₁₇H₁₅N₃O₄S: C, 57.13; H, 4.23; N 11.75. Found: C, 57.21; H, 4.31; N 11.73.

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Example 12

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Methyl [5(6)-(4-hydroxyphenylthio)-1*H*-benzimidazol-2yl]carbamate from 4-(4-hydroxyphenylthio)-1,2-phenylenediamine (4g) and 1,3-bis(methoxycarbonyl)-S-methyl isothiourea (6g) there was obtained the title compound (2.4g) as a white solid

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15 m.p. 202–204°C. m/e 315 (M⁺). H-NMR (400 MHz, d6-DMSO) δ 3.74 (s, 3H₂), 6.76 and 7.2 (AA'BB', 4H₂, J = 8.6 Hz), 7.03 (dd, 1H₂, J = 1.7, 6.6 Hz₂), 7.29 (d, 1H₂, J = 1.4 Hz₂), 7.34 (d, 1H₂, J = 8.3 Hz₂), 9.68 (b, 1H₂), 11.67 (b, 2H) ppm.

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Example 13

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Methyl [5(6)-(4-(acetylamino)phenoxy)-1*H*-benzimidazol-2yl]carbamate from 4-(4-(acetylamino)phenoxy)-1,2-phenylenediamine (1.02g) and 1,3-bis(methoxycarbonyl)-S-methyl isothiourea (2.67g) there was obtained the title compound (0.93g) as a white solid m.p. 304-306°C m/e 340 (M⁺). Anal. Calculated for C₁₇H₁₆N₄O₄: C, 60.00; H, 4.74; N 16.46. Found: C, 60.03; H, 4.72; N 16.42.

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Prepared in an analogous fashion to Example 3 was:

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Example 14

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Methyl [5(6)-(4-phosphonooxyphenoxy)-1*H*-benzimidazol-2yl]carbamate from methyl [5(6)-(4-(di-*tert*-butylphosphonooxy)phenoxy)-1*H*-benzimidazol-2yl]carbamate

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(200mg) there was obtained the title compound (140mg) as a white solid m.p. 272-275°C.

Example 15

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Methyl [5(6)-(4-hydroxy-α-hydroxyiminobenzyl)-1*H*-benzimidazol-2yl]carbamate [5(6)-(4-hydroxybenzoyl)-1*H*-benzimidazol-2yl]carbamate (100 mg, 0.32 mmol) was added to a solution of hydroxylamine (0.8 mmol) in MeOH (20 mL, prepared by treatment of hydroxylamine hydrochloride (0.64 g, 1.6 mmol) and NaOH (0.14g, 1.6 mmol) followed by filtration). The solution was heated at 70°C for 36 h, cooled and water (30 ml) added. The solution was filtered and washed with ether and the solid triturated with methanol to yield the title compound (40 mg) as a white solid; mp 288–290°C.

The following known compounds were prepared by literature methods:

Example 16: Methyl [5(6)-(4-(acetylamino)phenoxy)-1H-benzimidazol-2yl]carbamate

Example 17: Methyl [5(6)-(4-aminophenoxy)-1H-benzimidazol-2yl]carbamate

20 Example 18: Methyl [5(6)-(3-aminophenoxy)-1H-benzimidazol-2yl]carbamate

Example 19: [5(6)-(4-hydroxybenzoyl)-1H-benzimidazol-2yl]carbamate

Example 20: [5(6)-(2-hydroxybenzoyl)-1H-benzimidazol-2yl]carbamate

Example 21: [5(6)-(3-hydroxybenzoyl)-1H-benzimidazol-2yl]carbamate

Claims

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CLAIMS

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1. Use of vascular damaging agents in the preparation of compositions for the treatment of diseases involving angiogenesis, characterised in that the agents are 5(6)-substituted benzimidazole-2-carbamates of formula:

I

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wherein

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Alk is an alkyl group

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X is oxygen, sulphur, sulphinyl, sulphonyl, carbonyl (CO), thiocarbonyl (CS), sulphonyloxy, NH, iminomethylene (C=NH), N-hydroxyiminomethylene, N-alkoxyiminomethylene, dialkoxymethylene, 1,3-dioxolan-2yl, 1,1-ethenyl, a group

15 CHR³ or a bond

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 R^1 is hydrogen, alkylaminocarbonyl or alkoxycarbonyl R^2 is hydrogen, alkoxycarbonyl, cyanomethyl, cyanoethyl, alkoxymethyl or acetoxymethyl.

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R3 is hydrogen, hydroxy, alkoxy or amino

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A is an optionally substituted aromatic, optionally substituted heteroaromatic, optionally substituted heterocycloalkyl, optionally substituted alkyl or optionally substituted cycloalkyl group

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and the pharmaceutically acceptable salts, solvates and hydrates thereof.

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2. A use according to claim 1 wherein the substituent on A is selected from (a) alkyl substituted by one or more of hydroxy, amino, alkylamino, dialkylamino, halogen, carboxyl, SO₃H, sulphate, phosphate, alkoxycarbonyl, aralkoxycarbonyl, alkoxycarbonylamino, aminoalkylaminocarbonyl and cyano, or halogen, hydroxy,

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amino, alkoxy, alkylthio, cyano, nitro, sulphate, isothiocyanate, aryl, heteroaryl or heterocycloalkyl; or (b) a group Y selected from phosphate, alkylphosphate, C(O)R⁴, OC(O)R⁴, SO₂R⁴, NHC(O)R⁴, NHC(O)R⁴, NR⁵C(O)R⁴, SR⁴, S(O)R⁴, OSO₂R⁴, NHSO₂R⁴,

NR⁵SO₂R⁴, SO₃H, CO₂H and CO₂R⁵ where R⁴ is selected from hydrogen, R⁵, OR⁵, NHR⁵, NR⁵R⁶, aryl, heteroaryl or heterocycloalkyl such aryl, heteroaryl or heterocycloalkyl groups being optionally substituted with one or more substituents selected from alkyl, heterocycloalkyl, haloalkyl, hydroxy, nitro, cyano, amino, alkylamino, dialkylamino, halogen, carboxyl, SO₃H, sulphate and phosphate wherein R⁵ and R⁶, which may be the same or different, are each an alkyl group optionally substituted with one or more substituents selected from hydroxy, amino, alkylamino, dialkylamino, guanidino, halogen, carboxyl, SO₃H, sulphate, phosphate, aryl and heteroaryl.

3. A 5(6)-substituted benzimidazole-2-carbamate of formula

I

wherein

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Alk is an alkyl group

CHR³ or a bond

X is oxygen, sulphur, sulphinyl, sulphonyl, carbonyl (CO), thiocarbonyl (CS), sulphonyloxy, NH, iminomethylene (C=NH), N-hydroxyiminomethylene, N-alkoxyiminomethylene, dialkoxymethylene, 1,3-dioxolan-2yl, 1,1-ethenyl, a group

 R^1 is hydrogen, alkylaminocarbonyl or alkoxycarbonyl R^2 is hydrogen, alkoxycarbonyl, cyanoethyl, cyanoethyl, alkoxymethyl or acetoxymethyl.

R3 is hydrogen, hydroxy, alkoxy or amino

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A is an optionally substituted aromatic, optionally substituted heteroaromatic, optionally substituted heterocycloalkyl, optionally substituted alkyl or optionally substituted cycloalkyl group

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in which at least one substituent is Y selected from phosphate, alkylphosphate, $C(O)R^4$, $OC(O)R^4$, SO_2R^4 , $NHC(O)R^4$, $NR^5C(O)R^4$, SR^4 , $S(O)R^4$, OSO_2R^4 , $NHSO_2R^4$, $NR^5SO_2R^4$, SO_3H , CO_2H and CO_2R^5 where R^4 is selected from hydrogen, R^5 , OR^5 , NHR^5 , NR^5R^6 , aryl, heteroaryl or heterocycloalkyl such aryl, heteroaryl or heterocycloalkyl groups being optionally substituted with one or more substituents selected from alkyl, heterocycloalkyl, haloalkyl, hydroxy, nitro, cyano, amino, alkylamino, dialkylamino, halogen, carboxyl, SO_3H , sulphate and phosphate wherein R^5 and R^6 , which may be the same or different, are each an alkyl group optionally substituted with one or more substituents selected from hydroxy, amino, alkylamino, dialkylamino, guanidino, halogen, carboxyl, SO_3H , sulphate, phosphate, aryl and heteroaryl and the pharmaceutically acceptable salts, solvates and hydrates thereof.

4. A carbamate according to claim 3 of formula:

20 wherein

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alk is an alkyl group

B is an aromatic or heteroaromatic ring

X is oxygen, sulphur, sulphinyl, sulphonyl, carbonyl (CO), thiocarbonyl (CS),

sulphonyloxy, NH, iminomethylene (C=NH), N-hydroxyiminomethylene, Nalkoxyiminomethylene, dialkoxymethylene, 1,3-dioxolan-2yl, 1,1-ethenyl, a group CHR³ or a bond

R¹ is hydrogen, alkylaminocarbonyl or alkoxycarbonyl

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10	R ² is hydrogen, alkoxycarbonyl, cyanomethyl, cyanoethyl, alkoxymethyl or acetoxymethyl R ³ is hydrogen, hydroxy, alkoxy or amino Y is as hereinbefore defined
15	R ⁷ and R ⁸ are each independently H, alkyl, halogen, hydroxy, amino, alkylamino, dialkylamino, alkoxy, alkylthio, cyano, nitro, or trifluoromethyl
20	with the proviso that Y is not NHC(O)Me and when B is a thiophene ring then Y is not C(O)CF ₃ and when B is a 5(6)-benzimidazole ring then Y is not NHCO ₂ Me or NHCO ₂ Et
25	and the pharmaceutically acceptable salts, solvates, hydrates and prodrugs thereof. 5. A carbamate according to claim 4 in which Y is a phosphate group.
30	 A carbamate according to claim 4 in which Y is NHC(O)R⁴ wherein R⁴ is a 1-aminoalkyl group.
35	 A composition for treatment of diseases involving angiogenesis comprising at least one 5(6)-substituted benzimidazole-2-carbamate of formula I as defined in claim 1 in an amount sufficient to damage new vasculature.
40	 A composition according to claim 7, wherein the carbamate is as defined in claim 2.

9. A composition according to claim 7, wherein the carbamate has a formula II as defined in claim 3.

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(54) Title: BENZIMIDAZOLE VASCULAR DAMAGING AGENTS

(54) Titre: AGENTS DE DEGRADATION VASCULAIRE AUX BENZIMIDAZOLES

(57) Abstract

5(6)-substituted benzimidazole-2-carbamates of formula (I) wherein Alk is an alkyl group, X is oxygen, sulphur, sulphur,

(57) Abrégé

La présente invention concerne un groupe d'agents de dégradation vasculaire convenant à la péparation de médicaments destinés au traitement d'affections impliquant une néovascularisation. Ces molécules sont des benzimidazole-2-carbamates à substitution 5(6) représentés par la formule générale (I) ou certains de leurs sels, solvates et hydrates admis en pharmacie. Dans cette formule, "alk" est un groupe alkyle. X est oxygène, soufre, sulphinyle, sulphonyle, carbonyle (CO), thiocarbonyle (CS), sulphonyloxy, NH, iminométhylène (C=NH), N-hydroxyiminométhylène, N-alcoxyiminométhylène, dialcoxyméthylène, 1,3-dioxolane-2yle, 1,1-éthényle, un groupe CHR3¿ ou une liaison. R1¿ est hydrogène, alkylaminocarbonyle ou alcoxycarbonyle. R2¿ est hydrogène, alcoxycarbonyle, cyanométhyle, cyanoéthyle, alcoxyméthyle ou acétoxyméthyle. R3¿ est hydrogène, hydroxy, alcoxy ou amino. A est un groupe éventuellement substitué, notamment aromatique, hétéroaromatique, hétérocycloalkyle, alkyle ou cycloalkyle. La plupart des composés de ce groupe entrent dans le cadre de l'invention, en particulier ceux dans lesquels A est un cycle aromatique ou hétéroaromatique à substituants, particulièrement des substituants qui sont phosphates ou alkylphosphates. L'invention concerne donc aussi bien les composés de l'invention que les compositions pharmaceutiques comportant des composés ressortissants de la définition au sens large.



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- (74) Agents: BAILLIE, lain, C. et al.; Languer Parry, 52-54 High Holborn, London WCIV 6RR (GB).

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(54) Title: BENZIMIDAZOLE VASCULAR DAMAGING AGENTS

(57) Abstract

5(6)-substituted benzimidazole-2-carbamates of formula (I) wherein Alk is an alkyl group, X is oxygen, sulphur, sulphinyl, sulphonyl, carbonyl (CO), thiocarbonyl (CS), sulphonyloxy, NH, iminomethylene (C=NH), N-hydroxyiminomethylene, Nalkoxyiminomethylene, dialkoxymethylene, 1,3-dioxolan-2yl, 1,1-ethenyl, a group CHR3 or a bond, R1 is hydrogen, alkylaminocarbonyl or alkoxycarbonyl, R2 is hydrogen, alkoxycarbonyl, cyanomethyl, cyanoethyl, alkoxymethyl or acetoxymethyl. R3 is hydrogen, hydroxy, alkoxy or amino. A is an optionally substituted aromatic, optionally substituted heteroaromatic, optionally substituted heterocycloalkyl, optionally substituted alkyl or optionally substituted cycloalkyl group and the pharmaceutically acceptable salt, solvates and hydrates thereof, can be used in the preparation of medicaments for the treatment of diseases involving neovascularisation.

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EE	Estonia	LR	Liberia	SG			
_				50	Singapore		

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A CLASS IPC 7	IFICATION OF SUBJECT MATTER A61K31/4184 A61P9/00 C07D23	5/32 C07F9/6506	
	o International Patent Classification (IPC) or to both national classi	fication and IPC	
	SEARCHED		
IPC 7	ocumentation searched (classification system followed by classific A61K A61P C07D C07F		
	tion searched other than minimum documentation to the extent tha	·	
	ata base consulted during the international search (name of data i		ed)
EPU-In	ternal, WPI Data, BEILSTEIN Data, (CHEM ABS Data	
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT		
Category *	Citation of document, with indication, where appropriate, of the r	olovant passages	Relevant to claim No.
X	KRUSE L I ET AL: "Synthesis, tubinding, antineoplastic evaluatistructure-activity relationship oncodazole analogues" JOURNAL OF MEDICINAL CHEMISTRY, vol. 32, no. 2, February 1989 (1 pages 409-17, XP002142971 the whole document, also page 41 I, compound 28	on, and of 989-02),	1-9
X Furth	er documents are listed in the continuation of box C.	Patent family members are listed	in annex
"A" documer consider a filing de filing de documer which is citation "O" documer other n" "P" documer later the Date of the a	ocument but published on or after the international atterms the published on or after the international atterms to other the publication date of another or other special reason (as specified) in terfering to an oral disclosure, use, exhibition or season to published prior to the international filing date but an the priority date claimed octual completion of the international search	"I" later document published after the inter or priority date and not in conflict with ofted to understand the principle or the invention." X' document of particular relevance; the carnot be considered novel or cannot involve an inventive step when the do. "Y" document of particular relevance; the carnot be considered to involve an interest is combined with one or no ments, such combination being obvious in the sit. "&" document member of the same patent if Date of mailing of the international sea.	the application but sony underlying the taimed invention be considered to current is taken alone fairmed invention rentive step when the re other such docu- us to a person skilled family
	July 2000	07/08/2000	
Name and m	alling address of the ISA European Patent Office, P.B. 5918 Patentham 2 NL - 2290 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nt,	Authorized officer Alland M	

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Inter inel Application No PCT/GB 00/00099

Category *	Lation) DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages	[Balance 11 11
		Relevant to claim No.
X	LACEY E ET AL: "Activity of benzimidazole carbamates against L1210 mouse leukaemia cells: Correlation with in vitro tubulin polymerization assay" BIOCHEMICAL PHARMACOLOGY, vol. 34, no. 19, 1985, pages 3603-3605, XP002056333 ISSN: 0006-2952 the whole document	1,2,7,8
x	CHEMICAL ABSTRACTS, vol. 84, no. 21, 24 May 1976 (1976-05-24) Columbus, Ohio, US; abstract no. 144640r, DE BRABANDER M J ET AL: "The effects of methyl '5-(2-thienylcarbonyl)-1H-benzimidazol-2-yl!carbamate, (R 17934; NSC 238159), a new synthetic antitumoral drug interfering with microtubules, on mammalian cells cultured in vitro" page 26; XP002143095 cited in the application abstract & CANCER RES., vol. 36, no. 3, 1976, pages 905-16,	1,2,7,8
X	US 3 965 113 A (BEARD C C ET AL) 22 June 1976 (1976-06-22) the whole document	3,4,7-9
X	US 3 694 455 A (DUNN G L) 26 September 1972 (1972-09-26) the whole document	3,7-9
X	DE 23 48 104 A (FARBWERKE HOECHST AG) 3 April 1975 (1975-04-03) the whole document, particularly compounds VIII	3
X	DE 21 64 690 A (FARBWERKE HOECHST AG) 12 July 1973 (1973-07-12) the whole document, particularly example 19	3,4,7-9
(DE 23 32 343 A (FARBWERKE HOECHST AG) 16 January 1975 (1975-01-16) the whole document, particularly example 20	3,4,7-9
	-/	

tnter mail Application No PCT/GB 00/00099

(Continu	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	
ategory *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	KHAN A M ET AL: "Sudies on enteric anthelmintics: Impact of single point structural change on the activity profile" ZEITSCHRIFT FÜR NATURFORSCHUNG, B, vol. 43, no. 2, February 1988 (1988-02), pages 233-7, XP002142972 the whole document	3,4,7-9
(RAEYMAEKERS A H M ET AL: "Synthesis and anthelmintic activity of alkyl-(5-acyl-1H-benzimidazol-2-yl) carbamates" ARZNEIMITTEL-FORSCHUNG, vol. 28(I), no. 4, 1978, pages 586-94, XP002142973 the whole document, particularly page 591, table 5, 22nd entry	3,4,7-9
X	ABUZAR S ET AL: "Synthesis and anthelmintic activity of 2,2'-disubstituted 5,5'-dibenzimidazolylsulfides and sulfones" ARZNEIMITTEL-FORSCHUNG, vol. 36(I), no. 3, March 1986 (1986-03), pages 416-9, XP002142974 the whole document	3,7-9
X	DATABASE CROSSFIRE 'Online! Beilstein Institut zur Foerderung der Chemischen Wissenschaften; XP002142975 Beilstein Registry Number 4595458 & INDIAN J. CHEM. SECT. B, vol. 19, no. 7, 1980, pages 536-8,	3,4
(DATABASE CROSSFIRE 'Online! Beilstein Institut zur Foerderung der Chemischen Wissenschaften; XP002142976 Beilstein Registry Number 5651135 & INDIAN J. CHEM. SECT. B, vol. 24, 1985, pages 754-60,	3,4,7-9
	DATABASE CROSSFIRE 'Online! Beilstein Institut zur Foerderung der Chemischen Wissenschaften; XP002142977 Beilstein Registry Number 5622648 & INDIAN J. CHEM. SECT. B, vol. 24, 1985, pages 747-53,	3,4,7-9

Interr viel Application No PCT/GB 00/00099

C.(Continu	ntion) DOCUMENTS CONSIDERED TO BE RELEVANT	PC1/68 00/00099			
Category *		Relevant to claim No.			
X	DATABASE CROSSFIRE 'Online! Beilstein Institut zur Foerderung der Chemischen Wissenschaften; XP002142978 Beilstein Registry Number 5669479, 5670801, 5667941, 5667940, 5665325 å INDIAN J. CHEM. SECT. B, vol. 24, 1985, pages 730-2,	3,7-9			
X	DATABASE CROSSFIRE 'Online! Beilstein Institut zur Foerderung der Chemischen Wissenschaften; XP002142979 Beilstein Registry Number 4615289, 4609187 & INDIAN J. CHEM. SECT. B, vol. 23, no. 12, 1984, pages 1274-8,	3,7-9			
X	DATABASE CROSSFIRE 'Online! Beilstein Institut zur Foerderung der Chemischen Wissenschaften; XP002142980 Beilstein Registry Number 6009722, 6007941, 6007769 & INDIAN J. CHEM. SECT. B, vol. 24, 1985, pages 178-81,	3,7-9			
A ·	US 5 763 473 A (ELOKDAH H M ET AL) 9 June 1998 (1998-06-09) the whole document	1			
	÷				

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

Continuation of Box I.2

Claims Nos.: 1-4, 7-9 (all partly)

The initial phase of the search revealed a very large number of documents relevant to the issue of novelty of claims 1-4 and 7-9. So many documents were retrieved that it is impossible to determine which parts of these claims may be said to define subject-matter for which protection might legitimately be sought (Article 6 PCT). For these reasons, a meaningful search and an exhaustive seach report over the whole breadth of said claims are impossible. Consequently, the documents cited in the search report with regard to said claims should only be considered as forming a representative sample of the revealed documents.

domation on patent family members

Inter Inal Application No PCT/GB 00/00099

Patent document cited in search repo		Publication date		Patent family member(s)	Publication date
US 3965113	A	22-06-1976	AR	214701 A	
	7	EC 00-13/0	AK	332422 B	31-07-1979 27-09-1976
			AT	1075073 A	15 - 01-1976
			Ãΰ	6367573 A	19-06-1975
			CA	1023750 A	03-01-1978
			CH	592634 A	31-10-1978
			DE	2363348 A	18 - 07-1974
			DK	151628 B	21-12-1987
			ES	421928 A	16-10-1976
			FR	2212149 A	26-07-1974
			GB	1434830 A	05-05-1976
			ĬĒ	40141 B	28-03-1979
			ĪĹ	43859 A	31-12-1976
			IN	138651 A	06-03-1976
			IN	138645 A	06-03-1976
			JP	49094671 A	09-09-1974
			NL	7317798 A	02-07-1974
			AR	216037 A	30-11-1979
			AR	221816 A	31-03-1981
			AT	341822 B	27-02-1978
			AT	835875 A	15-06-1977
			AT	332883 B	25-10-1976
			AT	1075173 A	15-02-1976
		,	AU	6367473 A	1 9- 06-1975
			BE	809234 A	28-06-1974
			BE	809235 A	28-06-1974
			CA	1032171 A	30-05-1978
			CH	608006 A	15-12-1978
			CH	613955 A	31-10-1979
			CS	187390 B	31-01-1979
			CS	187395 B	31-01-1979
			DD	112450 A	12-04-1975
			DE	2363351 A	11-07-1974
			DE	2366069 A	10-11-1977
			DE	2366070 C	11-06-1987
			DK	137329 B	20-02-1978
			ES ES	421927 A	01-01-1977
			FR	445257 A 2212150 A	01-10-1977 26-07-1974
			F R F R	2212150 A 2272665 A	26-07-1974 26-12-1075
			F K GB	2272005 A 1456497 A	26-12-1975 24-11-1976
			6B	1455728 A	24-11-1976 17-11-1976
			HK	8280 A	14-03-1980
			HÜ	169272 B	28 - 10-1976
			ÏE	40046 B	28-02-1979
			ĬĹ	43860 A	31 - 05-1977
			IN	138650 A	06-03-1976
US 3694455	Α	26-09-1972	NONE	7	
DE 2348104	Α	03-04-1975	AU	7358674 A	25 02 1022
~_ EUTUIU 1	^	00 UT-13/0	BE	/3586/4 A 820324 A	25-03-1976 25-03-1075
			DD	820324 A 115493 A	25-03-1975 05-10-1975
			DK	115493 A 502574 A	05-10-1975 02-06-1975
			FI	276974 A	02-06-1975 26-03-1975
			FR	276974 A 2244504 A	20-03-1975 18-04-1975
			JP	50058072 A	18-04-1975 20-05-1975
			NL	7412464 A	20-05-1975 27-03-1975
					-

Jornation on patent family members

of notsoligga land PCT/GB 00/00099

			PCT/GB 00/00099				
Patent document cited in search report	Publication date		Patent family member(s)	Publication date			
DE 2348104 A		NO		21-04-1975			
		SE	7412041 A	26-03-1975			
		US	3928375 A	23-12-1975			
DE 2164690 A	12-07-1973	AT	320638 B	25-02-1975			
		AU	465226 B	18-09-1975			
		LUA	5048672 A	27-06-1974			
		BE	793358 A	27-06-1973			
		BG BG	22080 A	25-11-1976			
İ		CA	20591 A 1017750 A	05-12-1975 20-09-1977			
		CH	583710 A	14-01-1977			
1		CH	580080 A	30-09-1976			
		CS	173650 B	28-02-1977			
1		CS	173615 B	28-02-1977			
1		DD DD	109219 A 110760 A	20-10-1974			
•		DK	130293 B	12-01-1975			
		EG	11122 A	03-02-1975 31-05-1977			
		ES	409860 A	16-11-1975			
		FI	54299 B	31-07-1978			
		FR	2166048 A	10-08-1973			
		GB Hu	1360180 A 165060 B	17-07-1974			
		ΪΕ	37855 B	28-06-1974 26-10-1977			
		ĨĹ	41170 A	31-01-1977			
·		IT	1043888 B	29-02-1980			
		JP	50033067 B	27-10-1975			
		JP Ke	48080558 A 2613 A	29-10-1973			
		KR	7800116 A	15-04-1976 15-04-1978			
		MX	3541 E	10-02-1981			
		MY	9276 A	31-12-1976			
		NL	7217532 A,B	29-06-1973			
		NO Ph	137092 B 10285 A	19-09-1977			
		RO	71503 A	05-11-1976 30-01-1981			
İ		SE	395890 B	29-08-1977			
		SU	492086 A	15-11-1975			
		SU	493967 A	28-11-1975			
İ		US US	3954791 A 3984561 A	04-05-1976			
		ΥŬ	177773 A	05-10-1976 13-11-1981			
		ZA	7209076 A	28-11-1973			
DE 2332343 A	16-01-1975	AT	342041 B	10-03-1978			
		AT	525274 A	15-07-1977			
		CA CH	1031781 A	23-05-1978			
		DK	614706 A 341474 A	14 - 12-1979 17 - 03-1975			
		ES	427455 A	16-07-1976			
		HU	168527 B	28-05-1976			
		IT	1056735 B	20-02-1982			
,		JP JP	1163768 C	26-08-1983			
		JP	50040564 A 57058343 B	14-04-1975 09-12-1982			
		NL	7408394 A	30-12-1974			
		SE	7408321 A	27-12-1974			
Form PCT/ISA/210 (patent family ennex) (July 1992)							

commation on patent family members

PCT/GB 00/00099

			bera		PCT/GB 00/00099		
Patent document cited in search report	t	Publication date	Patr me	ent family ember(s)		Publication date	
DE 2332343	A		YU	1780	74 A	18-06-1982	
US 5763473	A	09-06-1998	NONE				
							
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						•	
						•	
					*		